AFNI & FMRI

Introduction, Concepts, Principles











http://afni.nimh.nih.gov/afni

Clinical uses are not supported or advised.

AFNI User

AFNI = Analysis of Functional Neurolmages

- Developed to provide an environment for FMRI data analyses
 - ★ And a platform for development of new software
- **AFNI** refers to both the program of that name and the entire package of external programs and plugins (more than 100)
- Important principles in the development of AFNI:
 - ★ Allow user to stay close to the data and view it in many different ways
 - ★ Give users the power to assemble pieces in different ways to make customized analyses
 - "With great power comes great responsibility"
 - to understand the analyses and tools
 - ★ "Provide mechanism, not policy"
 - ★ Allow other programmers to add features that can interact with the rest of the package

Principles (and Caveats) We* Live By

- Fix significant bugs as soon as possible
 - ★ But, we define "significant"
- Nothing is secret or hidden (AFNI is open source)
 - ★ <u>But</u>, possibly not very well documented or advertised
- Release early and often
 - * All users are beta-testers for life
- Help the user (message board; consulting with NIH users)
 - ★ Until our patience expires
- Try to anticipate users' future needs
 - ★ What we think you will need may not be what you actually end up needing

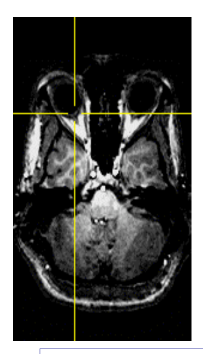
Outline of This Talk

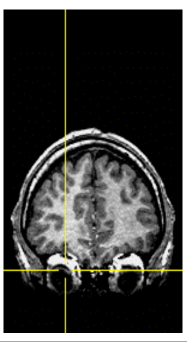
- Quick introduction to FMRI physics and physiology
 - ★ So you have some idea of what is going on in the scanner and what is actually being measured
- Brief discussion of FMRI experimental designs
 - ★ Block, Event-Related, Hybrid Event-Block
 - ★ But this is not a course in how to design your FMRI experimental paradigm
- Outlines of standard FMRI processing pipeline (AFNI-ized)
 - ★ Keep this in mind for the rest of the class!
 - ★ Many experiments require tweaking this "standard" collection of steps to fit the design of the paradigm and/or the inferential goals
- Overview of basic AFNI concepts
 - ⋆ Datasets and file formats; Realtime input; Controller panels; SUMA; Batch programs and Plugins

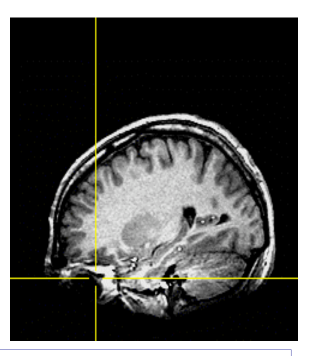
Quick Intro to MRI and FMRI

Physics and Physiology

(in pretty small doses)







MRI = Cool (and useful) Pictures

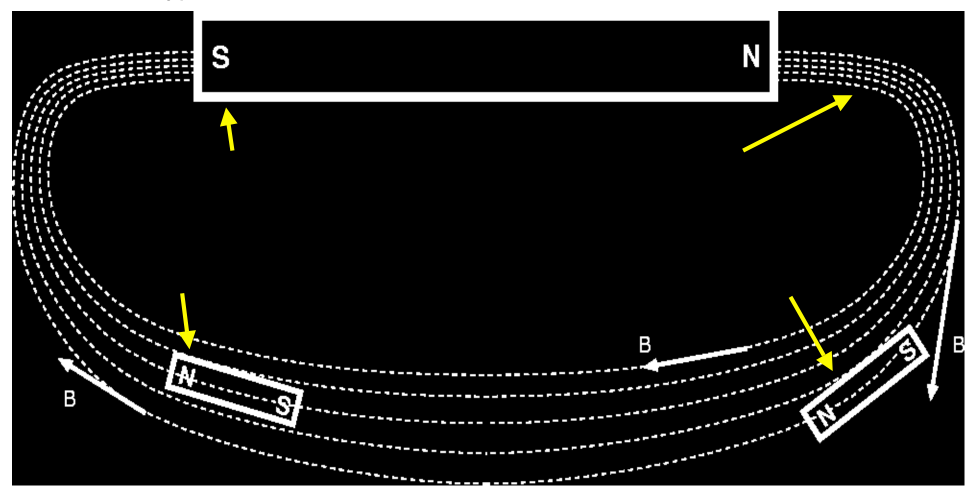
2D slices extracted from a 3D (volumetric) image [resolution about 1×1×1 mm; acquisition time about 10 min]

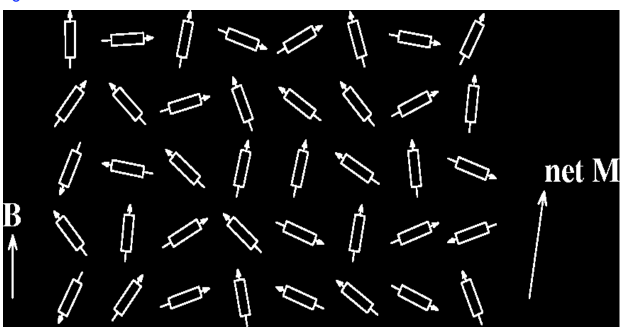
Synopsis of MRI

- 1) Put subject in big magnetic field (leave him there)
- 2) Transmit radio waves into subject [about 3 ms]
- 3) Turn off radio wave transmitter
- 4) Receive radio waves re-transmitted by subject
 - ★ Manipulate re-transmission with magnetic fields during this *readout* interval [10-100 ms]
- 5) Store measured radio wave data vs. time
 - ★ Now go back to 2) to get some more data
- 6) Process raw data to reconstruct images
- 7) Allow subject to leave scanner (optional)
- 8) Process images to extract desired features

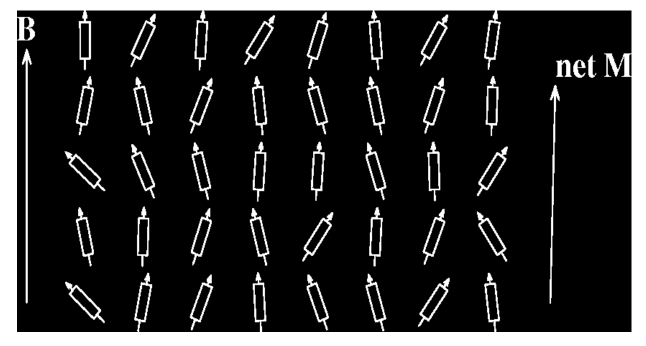
B_0 = Big Field Produced by Main Magnet

- Purpose is to align H protons in H₂O (little magnets)
- Units of B are Tesla (Earth's field is about 0.00005 Tesla)
 - ★ Typical field used in FMRI is 3 Tesla





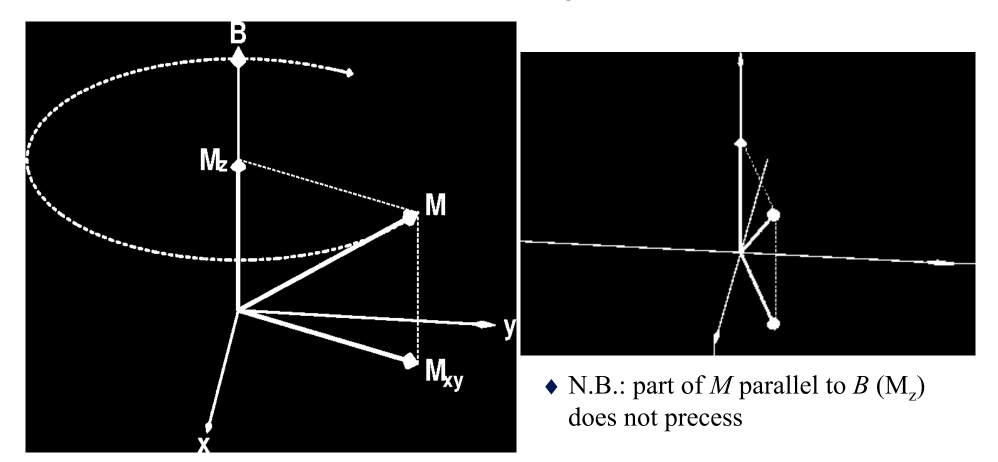
- ♦ Subject is <u>magnetized</u>
- Small B_0 produces small net magnetization M
- ♦ Thermal energy tries to randomize alignment of proton magnets



- Larger B_0 produces larger net magnetization M, lined up with B_0
- Reality check:
 0.0003% of protons aligned per Tesla of B₀

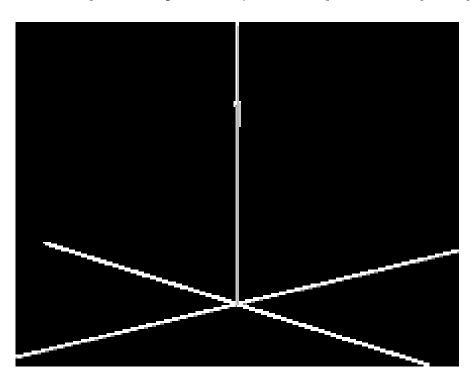
Precession of Magnetization M

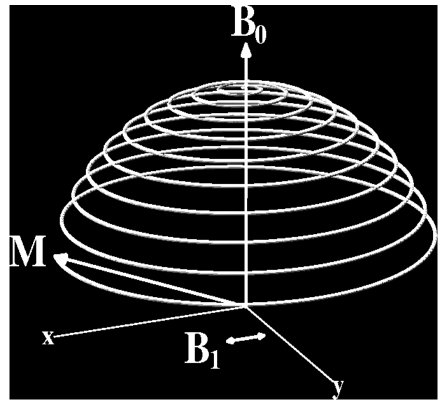
- Magnetic field B causes M to rotate ("precess") about the direction of B at a frequency proportional to the size of B 42 million times per second (42 MHz), per Tesla of B
 - \star 127 MHz at B = 3 Tesla range of radio frequencies



B_1 = Excitation (Transmitted) RF Field

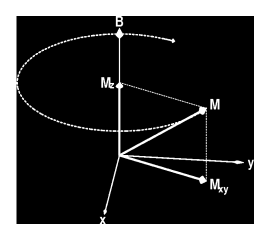
- Left alone, M will align itself with B in about 2–3 s
 - ⋆ ⇒ No precession ⇒ no detectable signal
- So don't leave it alone: apply (transmit) a magnetic field B₁ that fluctuates at the precession frequency (radio frequency=RF) and points perpendicular to B₀





Readout RF

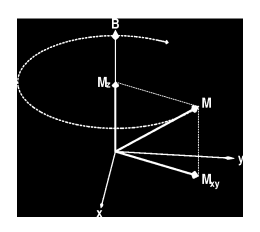
• When excitation RF is turned off, M is left pointed off at some angle to B_0 [flip angle]



- Precessing part of M [M_{xy}] is like having a magnet rotating around at very high speed (at RF speed: millions of revs/second)
- Will generate an oscillating voltage in a coil of wires placed around the subject — this is magnetic *induction*
- This voltage is the RF signal = the raw data for MRI
 - At each instant t, can measure one voltage V(t), which is proportional to the sum of all transverse M_{xy} inside the coil
 - Must separate signals originating from different regions
 - By reading out data for 5-60 ms, manipulating B field, being clever ...
 - Then have **image** of M_{xy} = map of how much signal from each *voxel*

Relaxation: Nothing Lasts Forever

- In the absence of external B_1 , M will go back to being aligned with static field $B_0 = relaxation$
- Part of M perpendicular to B₀ shrinks [M_{xy}]
 - \star This part of M = transverse magnetization
 - ★ It generates the detectable RF signal
 - ★ The relaxation of M_{xy} during readout affects the image
- Part of M parallel to B₀ grows back [M_z]
 - \star This part of M = longitudinal magnetization
 - * Not directly detectable, but is converted into transverse magnetization by external B_1
 - Therefore, M_z is the *ultimate* source of the NMR signal, but is not the *proximate* source of the signal



Time scale for this relaxation is called T2 or T2* = 20-40 ms in brain

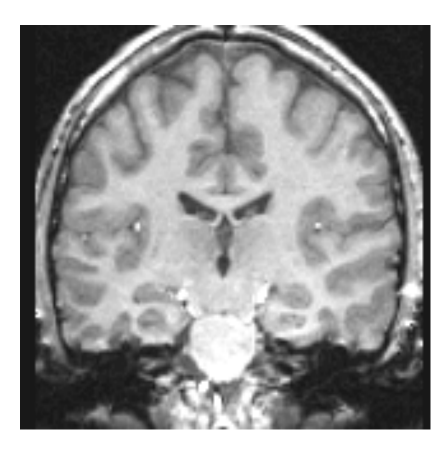
Time scale for this relaxation is called T1 = 500-2500 ms

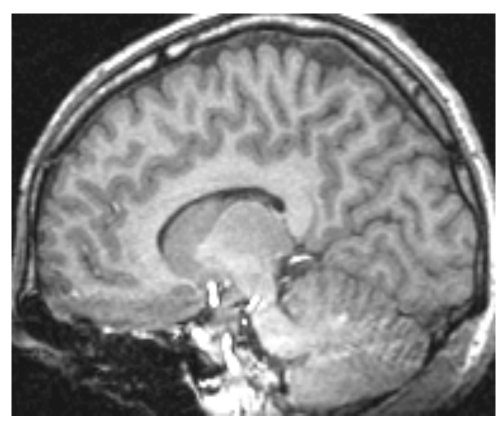
Material Induced Inhomogeneities in B

- Adding a nonuniform object (like a person) to B₀ will make the total magnetic field B nonuniform
 - ★ This is due to *susceptibility*: generation of extra magnetic fields in materials that are immersed in an external field
 - ⋆ Diamagnetic materials produce negative B fields [most tissue]
 - ★ Paramagnetic materials produce positive B fields [deoxyhemoglobin]
 - ★ Size of changes about $10^{-7} \cdot B_0 = 1 100$ Hz change in precession
- Makes the precession frequency nonuniform, which affects the image intensity and quality
 - ★ For large scale (100+ mm) inhomogeneities, scanner-supplied nonuniform magnetic fields can be adjusted to "even out" the ripples in B — this is called *shimming*
 - ⋆ Nonuniformities in B bigger than voxel size distort whole image
 - ★ Nonuniformities in B smaller than voxel size affect voxel "brightness"

The Concept of Contrast (or Weighting)

- Contrast = difference in RF signals emitted by water protons — between different tissues
- <u>Example</u>: gray-white contrast is possible because T1 is different between these two types of tissue





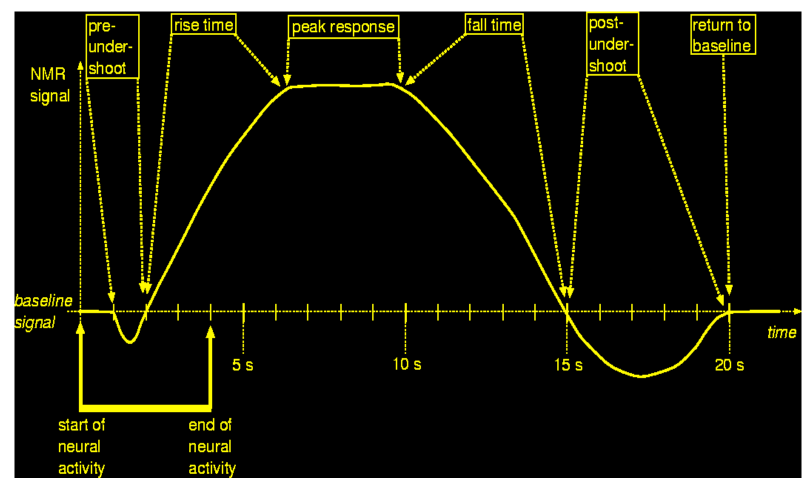
Types of Contrast Used in Brain FMRI

- T1 contrast at high spatial resolution
 - ★ Technique: use very short timing between RF shots (small TR) and use large flip angles
 - ★ Useful for anatomical reference scans
 - ★ 5-10 minutes to acquire 256×256×128 volume
 - ★ 1 mm resolution easily achievable
 - o finer voxels are possible, but acquisition time increases a lot
- T2 (spin-echo) and T2* (gradient-echo) contrast
 - ★ Useful for functional activation studies
 - \star 100 ms per 64×64 2D slice \Rightarrow 2-3 s to acquire whole brain
 - ★ 4 mm resolution
 - better is possible with better gradient system, and/or multiple RF readout coils

What is Functional MRI?

• 1991: Discovery that MRI-measurable signal increases a few % *locally* in the brain subsequent to increases in neuronal activity (Kwong, *et al.*)

Cartoon of MRI signal in an "activated" brain voxel



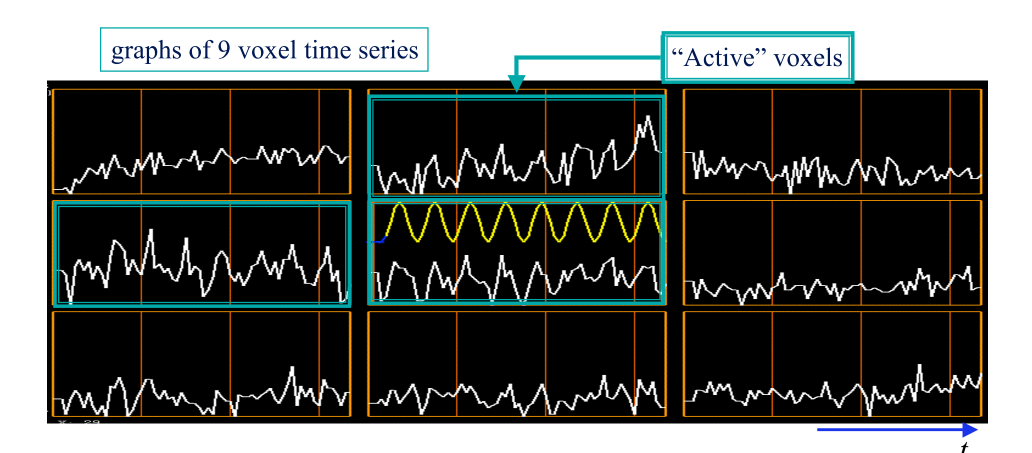
How FMRI Experiments Are Done

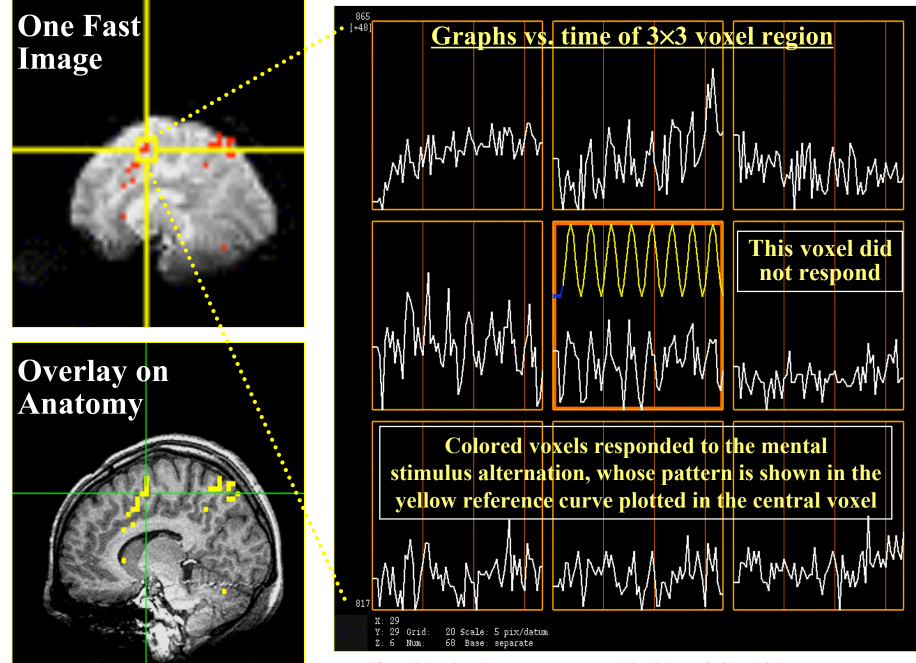
- Alternate subject's neural state between 2 (or more) conditions using sensory stimuli, tasks to perform, ...
 - ★ Can only measure relative signals, so must look for changes in the signal between the conditions
- Acquire MR images repeatedly during this process
- Search for voxels whose NMR signal time series (up-and-down) matches the stimulus time series pattern (on-and-off)
- Signal changes due to neural activity are small
 - Need 1000 or so images in time series (each slice) ⇒
 takes an hour or so to get reliable activation maps
 - Must break image acquisition into shorter "runs" to give the subject and scanner some break time
 - Other small effects can corrupt the results ⇒
 postprocess the data to reduce these effects & be careful
- Lengthy computations for image recon and temporal pattern matching

 data analysis usually done offline

Some Sample Data Time Series

- 16 slices, 64×64 matrix, 68 repetitions (TR=5 s)
- Task: phoneme discrimination: 20 s "on", 20 s "rest"





68 points in time 5 s apart; 16 slices of 64×64 images

Why (and How) Does NMR Signal Change With Neuronal Activity?

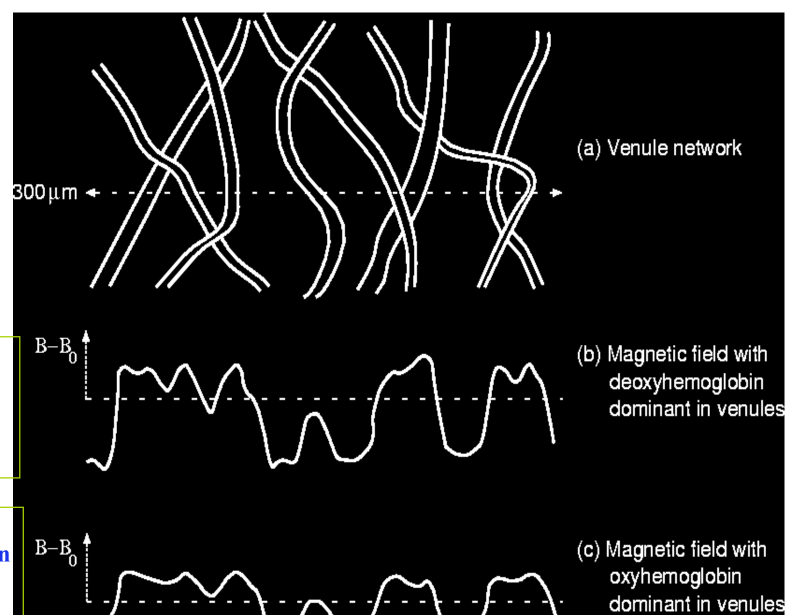
- There must be something that affects the water molecules and/or the magnetic field inside voxels that are "active"
 - ⋆ neural activity changes blood flow and oxygen usage
 - ★ blood flow changes which H₂O molecules are present and also changes the magnetic field locally
- FMRI is thus *doubly* indirect from physiology of interest (synaptic activity)
 - * also is much slower: 4-6 seconds after neurons
 - ★ also "smears out" neural activity: cannot resolve 10-100 ms timing of neural sequence of events

Neurophysiological Changes & FMRI

- There are 4 changes caused by neural activity that are currently observable using MRI:
- Increased Blood Flow
 - ★ New protons flow into slice from outside
 - ★ More protons are aligned with B₀
 - ★ Equivalent to a shorter T1 (as if protons are realigned faster)
 - ⋆ NMR signal goes up [mostly in arteries]
- Increased Blood Volume (due to increased flow)
 - ★ Total deoxyhemoglobin increases (as veins expand)
 - ★ Magnetic field randomness increases [more paramagnetic stuff in blood vessels]
 - ★ NMR signal goes down [near veins and capillaries]

- BUT: "Oversupply" of oxyhemoglobin after activation
 - ⋆ Total deoxyhemoglobin decreases
 - ★ Magnetic field randomness decreases [less paramag stuff]
 - ⋆ NMR signal goes up [near veins and capillaries]
 - ★ This is the important effect for FMRI as currently practiced
- Increased capillary perfusion
 - ★ Most inflowing water molecules exchange to parenchyma at capillaries
 - o i.e., the water that flows into a brain capillary is *not* the water that flows out!
 - ★ Can be detected with perfusion-weighted imaging methods
 - ★ This factoid is also the basis for ¹⁵O water-based PET
 - ★ May someday be important in FMRI, but is hard to do now

Cartoon of Veins inside a Voxel



Deoxyhemoglobin is paramagnetic (increases B)

Rest of tissue +oxyhemoglobin is diamagnetic (decreases B)

BOLD Contrast

- <u>BOLD</u> = <u>Blood Oxygenation Level Dependent</u>
- Amount of deoxyhemoglobin in a voxel determines how inhomogeneous that voxel's magnetic field is at the scale of the blood vessels (and red blood cells)
- Increase in oxyhemoglobin in veins after neural activation means magnetic field becomes more uniform inside voxel
 - ★ So NMR signal goes up (T2 and T2* are larger), since it doesn't decay as much during data readout interval
 - ★ So MR image is brighter during "activation" (a little)

• **Summary**:

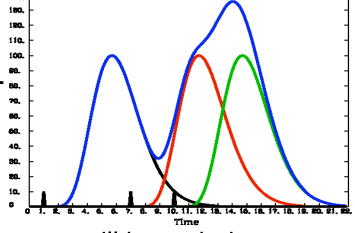
- ★ NMR signal increases 4-6 s after "activation", due to hemodynamic (blood) response
- ★ Increase is same size as noise, so need lots of data

FMRI Experiment Design and Analysis

- FMRI experiment design
 - **★ Event-related, block, hybrid event-block?**
 - ⋆ How many types of stimuli? How many of each type? Timing (intra- & inter-stim)?
 - ★ Will experiment show what you are looking for? (Hint: bench tests)
 - ★ How many subjects do you need? (Hint: the answer does not have 1 digit)
- Time series data analysis (individual subjects)
 - ★ Assembly of images into AFNI datasets; Visual & automated checks for bad data
 - ★ Registration of time series images
 - ⋆ Smoothing & masking of images; Baseline normalization; Censoring bad data
 - **★** Catenation into one big dataset
 - ★ Fit statistical model of stimulus timing+hemodynamic response to time series data
 - Fixed-shape or variable-shape response models
 - **★** Segregation into differentially active blobs
 - Thresholding on statistic + clustering <u>and/or</u> Anatomically-defined ROI analysis
 - ★ Visual examination of maps and fitted time series for validity and meaning
- Group analysis (inter-subject)
 - ★ Spatial normalization to Talairach-Tournoux atlas (or something like it)
 - **★** Smoothing of fitted parameters
 - Automatic global smoothing + voxel-wise analysis or ROI averaging
 - * ANOVA to combine and contrast activation magnitudes from the various subjects
 - ⋆ Visual examination of results (usually followed by confusion)
 - ★ Write paper, argue w/ mentor, submit paper, argue w/ referees, publish paper, ...

FMRI Experiment Design - 1

- Recall hemodynamic (FMRI) response
 - ⋆ peak is 4-6 s after neural activation
 - ★ width is 4-5 s for very brief (< 1 s) activation



- ★ ⇒ two separate activations less than 12-15 s apart will have their responses overlap and add up (approximately — more on this in a later talk!)
- Block design experiments: Extended activation, or multiple closely-spaced (< 2-3 s) activations
 - ★Multiple FMRI responses overlap and add up to something more impressive than a single brief blip
 - <u>★But</u> can't distinguish distinct but closely-spaced activations; example:
 - Each brief activation is "subject sees a face for 1 s, presses button #1 if male, #2 if female" and faces come in every 2 s for a 20 s block, then 20 s of "rest", then a new faces block, etc.
 - o What to do about trials where the subject makes a mistake? These are presumably neurally different than correct trials, but there is no way to separate out the activations when the hemodynamics blurs so much in time.

FMRI Experiment Design - 2

Event-related designs:

- ★ Separate activations in time so can model the FMRI response from each separately, as needed (e.g., in the case of subject mistakes)
- ★ Need to make inter-stimulus intervals vary ("jitter") if there is any potential time overlap in their FMRI response curves; e.g., if the events are closer than 12-15 s in time
 - Otherwise, the tail of event #x always overlaps the head of event #x+1 in the same way, and as a result the amplitude of the response in the tail of #x can't be told from the response in the head of #x+1

* Important note!

- You cannot treat every single event as a distinct entity whose response amplitude is to be calculated separately!
- You must still group events into classes, and assume that all events in the same class evoke the same response.
 - → Approximate rule: 25+ events per class (with emphasis on the '+')
- o There is just too much noise in FMRI to be able to get an accurate activation map from a single event!

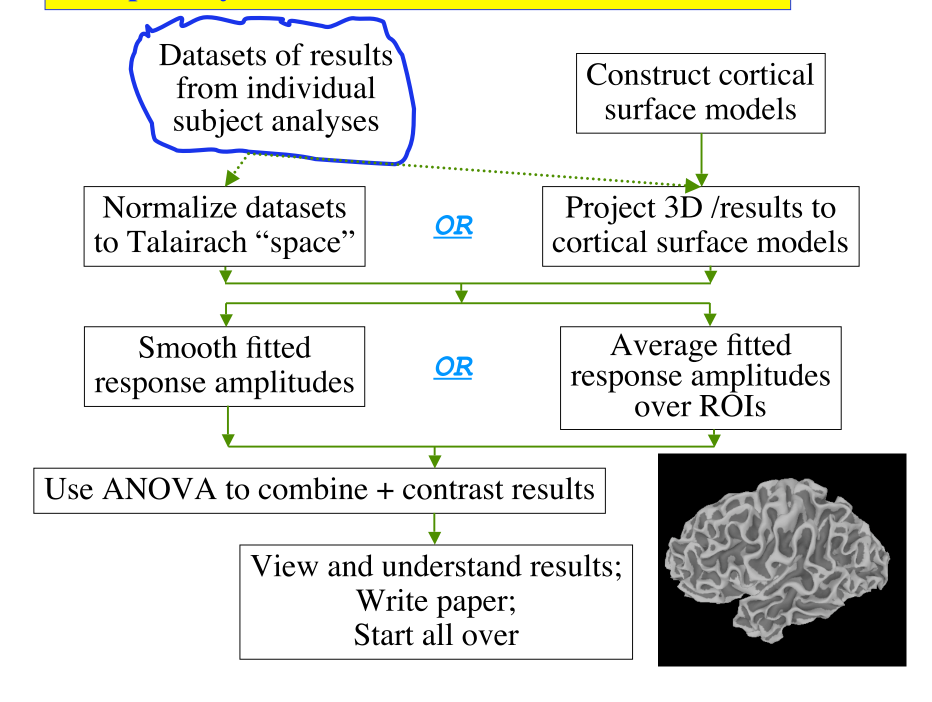
FMRI Experiment Design - 3

- Hybrid Block/Event-related designs:
 - ★ The long "blocks" are situations where you set up some continuing condition for the subject
 - ★ Within this condition, multiple distinct events are given
 - ★ Example:
 - Event stimulus is a picture of a face
 - Block condition is instruction on what the subject is to do when he sees the face:
 - → Condition A: press button #1 for male, #2 for female
 - → Condition B: press button #1 if face is angry, #2 if face is happy
 - o Event stimuli in the two conditions may be identical, or at least fungible
 - It is the instructional+attentional modulation between the two conditions that is the goal of such a study
 - → Perhaps you have two groups of subjects (patients and controls) which respond differently in bench tests
 - You want to find some neural substrates for these differences

3D Individual Subject Analysis

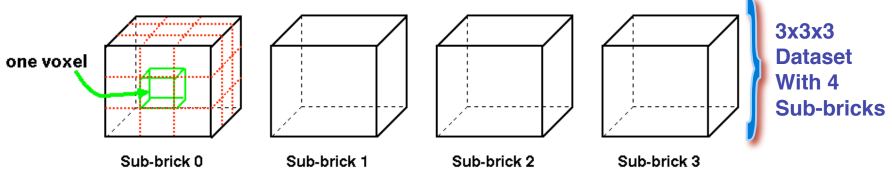
```
to3d
Assemble images into AFNI-formatted datasets
                                                     can do at scanner
Check images for quality (visual & automatic)
                              3dvolreg
Register (realign) images
                             3dWarpDrive
Smooth images spatially
                            3dmerge (optional)
Mask out non-brain parts of images
                                        3dAutomask + 3dcalc (optional)
Normalize time series baseline to 100 (for %-izing)
Fit stimulus timing + hemodynamic model to time series
   catenates imaging runs, removes residual movement
 effects, computes response sizes & inter-stim contrasts
                                                                3dDeconvolve
                                                    Alphasim + 3dmerge
Segregate into differentially "activated" blobs
                                                   Extraction from ROIs
                                   afni
Look at results, and think
                             your personal brain
.. to group analysis (next page)
```

Group Analysis: in 3D or on folded 2D cortex models



Fundamental AFNI Concepts

- Basic unit of data in AFNI is the <u>dataset</u>
 - ★ A collection of 1 or more 3D arrays of numbers
 - Each entry in the array is in a particular spatial location in a 3D grid (a <u>voxel</u> = 3D pixel)
 - Image datasets: each array holds a collection of slices from the scanner
 - ► Each number is the signal intensity for that particular voxel
 - Derived datasets: each number is computed from other dataset(s)
 - ⇒ e.g., each voxel value is a *t*-statistic reporting "activation" significance from an FMRI time series dataset, for that voxel
 - ★ Each 3D array in a dataset is called a sub-brick
 - There is one number in each voxel in each sub-brick

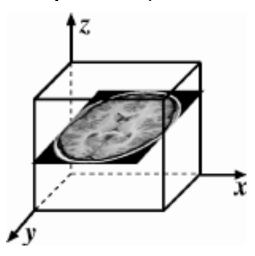


Dataset Contents: Numbers

- Different types of numbers can be stored in datasets
 - ★ 8 bit bytes (e.g., from grayscale photos)
 - ★ 16 bit short integers (e.g., from MRI scanners)
 - Each sub-brick may also have a floating point scale factor α attached, so that "true" value in each voxel is actually α (value in dataset file)
 - \star 32 bit floats (e.g., calculated values; lets you avoid the α)
 - ★ 24 bit RGB color triples (e.g., JPEGs from your digital camera!)
 - ★ 64 bit complex numbers (e.g., for the physicists in the room)
- Different sub-bricks are allowed to have different numeric types
 - ★ But this is *not* recommended
 - ★ Will occur if you "catenate" two dissimilar datasets together (e.g., using 3dTcat or 3dbucket commands)
 - Programs will display a warning to the screen if you try this

Dataset Contents: Header

- Besides the voxel numerical values, a dataset also contains auxiliary information, including (some of which is optional):
 - * xyz dimensions of each voxel (in mm)
 - ★ Orientation of dataset axes; for example, x-axis=R-L, y-axis=A-P, z-axis=I-S ⇒ axial slices (we call this orientation "RAI")
 - ★ Location of dataset in scanner coordinates
 - Needed to overlay one dataset onto another
 - Very important to get right in FMRI, since we deal with many datasets
 - ★ Time between sub-bricks, for <u>3D+time</u> datasets
 - Such datasets are the basic unit of FMRI data (one per imaging run)
 - ★ Statistical parameters associated with each sub-brick
 - o e.g., a t-statistic sub-brick has degrees-of-freedom parameter stored
 - o e.g., an F-statistic sub-brick has 2 DOF parameters stored



<u>AFNI Dataset Files - I</u>

- AFNI formatted datasets are stored in 2 files
 - ★ The <u>.HEAD</u> file holds all the auxiliary information
 - * The .BRIK file holds all the numbers in all the sub-bricks
- Datasets can be in one of 3 coordinate systems (AKA <u>views</u>)
 - ★ Original data or <u>+orig</u> view: from the scanner
 - ★ AC-PC aligned or <u>+acpc</u> view:
 - o Dataset rotated/shifted so that the anterior commissure and posterior commissure are horizontal (y-axis), the AC is at (x,y,z)=(0,0,0), and the hemispheric fissure is vertical (z-axis)
 - ★ Talairach or <u>+tlrc</u> view:
 - Dataset has also been rescaled to conform to the Talairach-Tournoux atlas dimensions (R-L=136 mm; A-P=172 mm; I-S=116 mm)
 - AKA <u>Talairach</u> or <u>Stererotaxic</u> coordinates
 - Not quite the same as MNI coordinates, but very close

AFNI Dataset Files - II

- AFNI dataset filenames consist of 3 parts
 - ★ The user-selected <u>prefix</u> (almost anything)
 - ★ The view (one of +orig, +acpc, or +tlrc)
 - ★ The <u>suffix</u> (one of .HEAD or .BRIK)
 - ★ Example: BillGates+tlrc.HEAD and BillGates+tlrc.BRIK
 - ★ When creating a dataset with an AFNI program, you supply the prefix; the program supplies the rest
- AFNI programs can *read* datasets stored in several formats
 - ★ ANALYZE (.hdr/.img file pairs); i.e., from SPM, FSL
 - ★ MINC-1 (.mnc); i.e., from mnitools
 - ★ CTF (.mri, .svl) MEG analysis volumes
 - ★ ASCII text (.1D) numbers arranged into columns
 - ★ Have conversion programs to write out MINC-1, ANALYZE, ASCII, and NIfTI-1.1 files from AFNI datasets, if desired

NIfTI Dataset Files

- NIfTI-1.1 (<u>.nii</u> or <u>.nii.gz</u>) is a new standard format that AFNI,
 SPM, FSL, BrainVoyager, et al., have agreed upon
 - ★ Adaptation and extension of the old ANALYZE 7.5 format
 - ★ Goal: easier interoperability of tools from various packages
- All data is stored in 1 file (cf. http://nifti.nimh.nih.gov/)
 - ★ 348 byte header (extensions allowed; AFNI uses this feature)
 - ★ Followed by the image numerical values
 - ★ Allows 1D-5D datasets of diverse numerical types
 - ★ .nii.gz suffix means file is compressed (with gzip)
- AFNI now reads and writes NIfTI-1.1 formatted datasets
 - ★ To write: when you give the prefix for the output filename, end it in ".nii" or ".nii.gz", and all AFNI programs will automatically write NIfTI-1.1 format instead of .HEAD/.BRIK
 - ★ To read: just give the full filename ending in ".nii" or ".nii.gz"

Dataset Directories

- Datasets are stored in directories, also called <u>sessions</u>
 - ★ All the datasets in the same session, in the same view, are presumed to be aligned in *xyz*-coordinates
 - o Voxels with same value of (x,y,z) correspond to same brain location
 - ★ Can overlay (in color) any one dataset on top of any other one dataset (in grayscale) from same session
 - Even if voxel sizes and orientations differ
 - ★ Typical AFNI contents of a session directory are all data derived from a single scanning session for one subject
 - Anatomical reference (T1-weighted SPGR or MP-RAGE volume)
 - 10-20 3D+time datasets from FMRI EPI functional runs
 - Statistical datasets computed from 3D+time datasets, showing activation (you hope and pray)
 - Datasets transformed from +orig to +tlrc coordinates, for comparison and conglomeration with datasets from other subjects

Getting and Installing AFNI

- AFNI runs on <u>Unix</u> systems: Linux, Sun, SGI, Mac OS X
 - ★ Can run under Windows with Cygwin Unix emulator
 - o This option is really just for trying it out not for production use
- If you are at the NIH: SSCC can install AFNI and update it on your system(s)
 - ★ You must give us an account with ssh access
- You can download precompiled binaries from our Website
 - ★ http://afni.nimh.nih.gov/afni
 - * Also: documentation, message board, humor, data, ...
- You can download source code and compile it
- AFNI is updated fairly frequently, so it is important to update occasionally
 - ★ We won't help you with old versions!

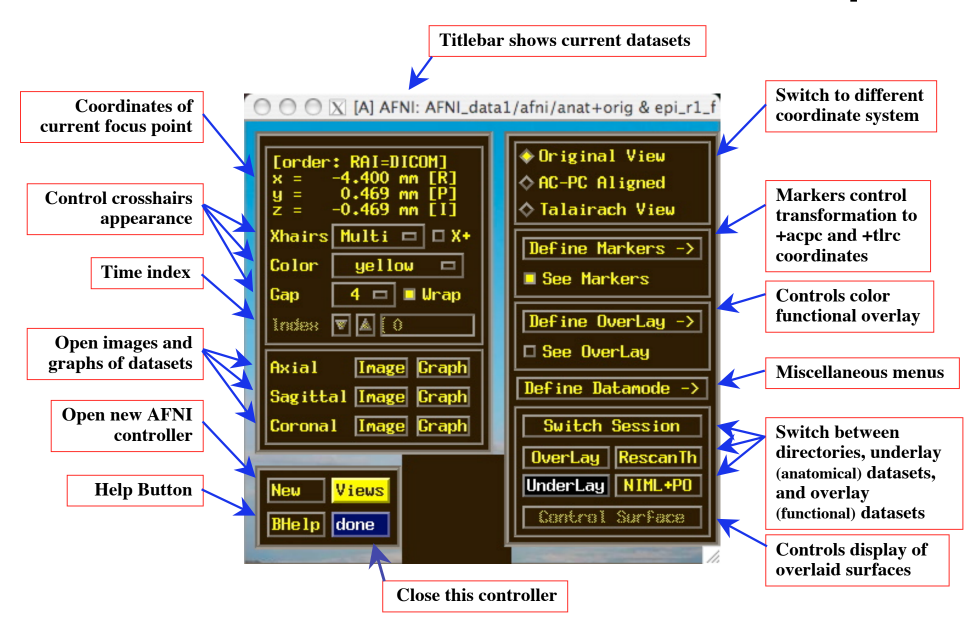
AFNI at the NIH Scanners

- AFNI can take images in "realtime" from an external program and assemble them into 3D+time datasets slice-byslice
- Jerzy Bodurka (FMRIF) has set up the GE Excite-based scanners (3T-1, 1.5 T, NMRF 3 T, and 7 T) to start AFNI automagically when scanning, and send reconstructed images over as soon as they are available:
 - ★ For immediate display (images and graphs of time series)
 - ★ Plus graphs of estimate subject head movement
- Goal is to let you see data as it is acquired, so that if there are any big problems, you can fix them right away
 - * Sample problem: someone typed in the imaging field-of-view (FOV) size wrong (240 cm instead of 24 cm), and got garbage data, *but only realized this too late* (after subject had left the scanner and gone home) *D'oh!*

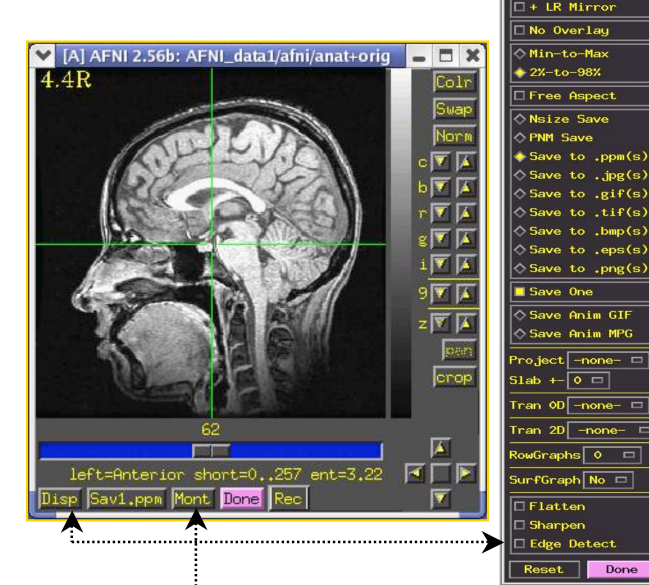
A Quick Overview of AFNI

- Starting AFNI from the Unix command line
 - ★ <u>afni</u> reads datasets from the current directory
 - * afni dir1 dir2 ... reads datasets from directories listed
 - ★ <u>afni -R</u> reads datasets from current directory and from all directories below it
- AFNI also reads a file named <u>.afnirc</u> from your home directory
 - ★ Used to change many of the defaults
 - Window layout and image/graph viewing setup; popup hints; whether to compress .BRIK files when writing
 - o cf. file README.environment in the AFNI documentation
- Also can read file <u>.afni.startup script</u> to restore the window layout from a previous run
 - ★ Created from Define Datamode->Misc->Save Layout menu
 - o cf. file README.driver for what can be done with AFNI scripts

AFNI controller window at startup



AFNI Image Viewer



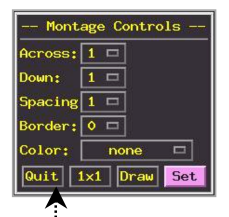
Disp and **Mont** control panels

No Rotation

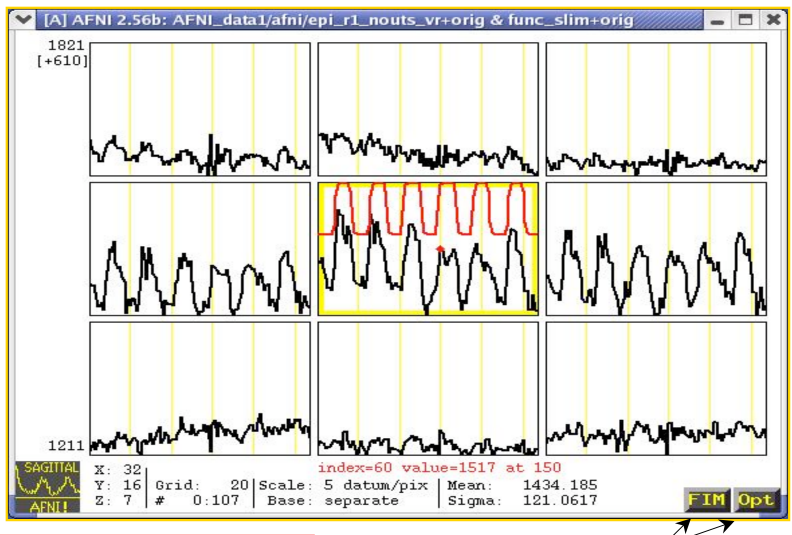
Done

CCW 90

♦ Rot 180 ♦ CW 90



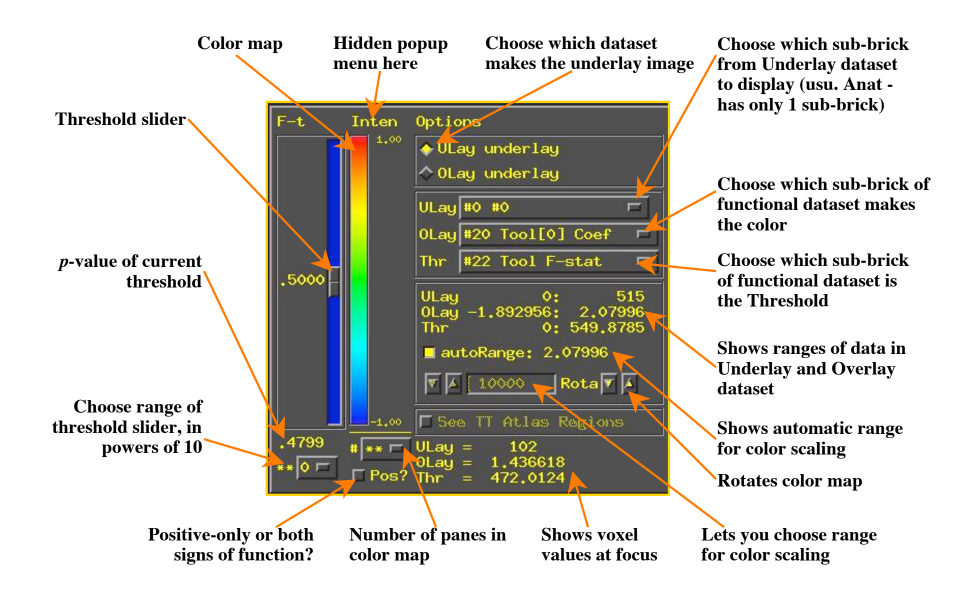
<u>AFNI Time Series Graph Viewer</u>



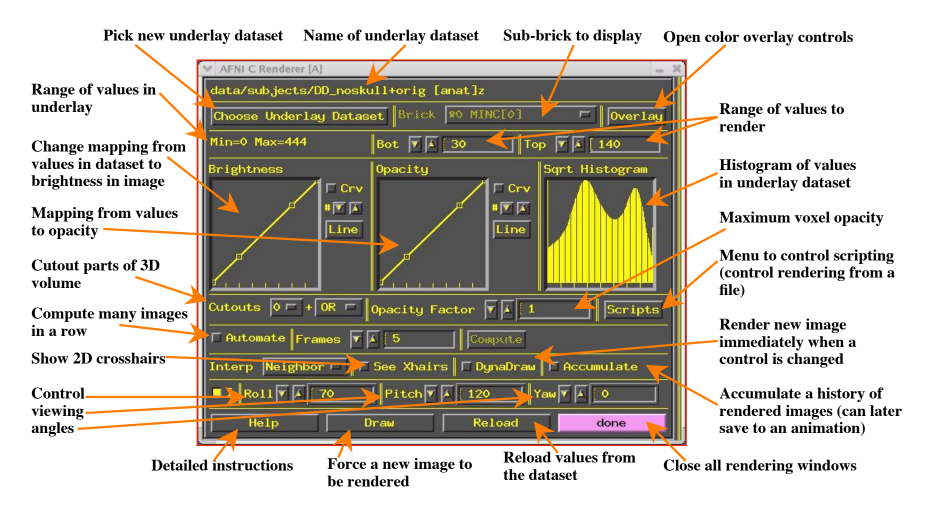
Data (black) and Reference waveforms (red)

Menus for controlling graph displays

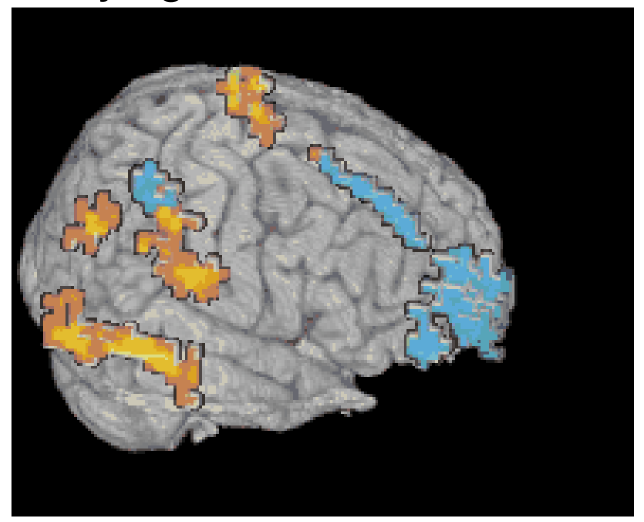
Define Overlay: Colorizing Panel (etc.)



Volume Rendering: an AFNI plugin



Staying Close to Your Data!



"ShowThru" rendering of functional activation: animation created with Automate and Save:aGif controls

Other Parts of AFNI

- Batch mode programs
 - ★ Are run by typing commands directly to computer, or by putting commands into a text file (script) and later executing them
- Good points about batch mode
 - ★ Can process new datasets exactly the same as old ones
 - ★ Can link together a sequence of programs to make a customized analysis (a personalized <u>pipeline</u>)
 - ⋆ Some analyses take a long time
- Bad points about batch mode
 - ★ Learning curve is "all at once" rather than gradual
 - ★ If you are, like, under age 35, you may not know how to, like, type commands into a computer
 - At least we don't make you use punched cards (yet)

AFNI Batch Programs

- Many important capabilities in AFNI are only available in batch programs
 - ★ A few examples (of more than 100, from trivial to complex)
- 3dDeconvolve = multiple linear regression on 3D+time datasets, to fit each voxel's time series to an activation model and then test these fits for significance
- 3dvolreg = 3D+time dataset registration, to correct for small subject head movements, and for inter-day head positioning
- <u>3dANOVA</u> = 1-, 2-, 3-, and 4- way ANOVA layouts, for combining & contrasting datasets in Talairach space
- <u>3dcalc</u> = general purpose voxel-wise calculator
- 3dclust = find clusters of activated voxels
- 3dresample = re-orient and/or re-size dataset voxel grid
- 3dSkullStrip = remove "skull" from anatomical dataset

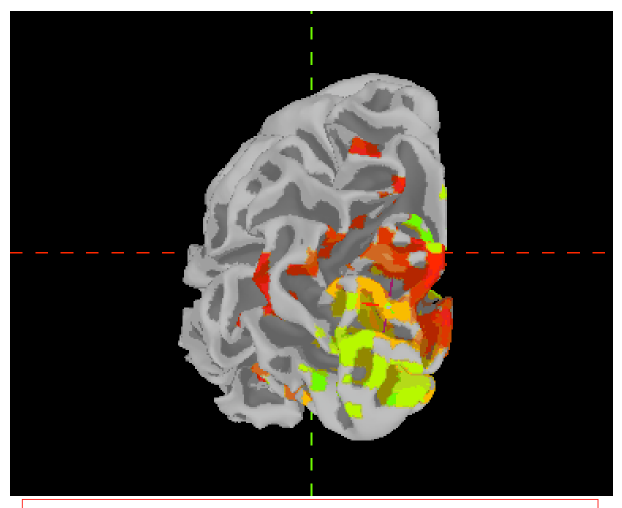
AFNI Plugins

- A <u>plugin</u> is an extension to AFNI that attaches itself to the interactive AFNI GUI
 - ★ Not the same as a batch program
 - ★ Offers a relatively easy way to add certain types of interactive functionality to AFNI
 - ★ A few examples:
- <u>Draw Dataset</u> = ROI drawing (draws numbers into voxels)
- Render [new] = Volume renderer
- <u>Dataset#N</u> = Lets you plot multiple 3D+time datasets as overlays in an AFNI graph viewer (e.g., fitted model over data)
- <u>Histogram</u> = Plots a histogram of a dataset or piece of one
- <u>Edit Tagset</u> = Lets you attach labeled "tag points" to a dataset (e.g., as anatomical reference markers)

SUMA, et alii

- <u>SUMA</u> is the AFNI <u>surface mapper</u>
 - ★ For displaying surface models of the cortex
 - Surface models come from <u>FreeSurfer</u> (MGH) or <u>SureFit/Caret</u> (Wash U) or <u>BrainVoyager</u>
 - ★ Can display functional activations mapped from 3D volumes to the cortical surface
 - ★ Can draw ROIs directly on the cortical surface o vs. AFNI: ROIs are drawn into the volume
- SUMA is a separate program from AFNI, but can "talk" to AFNI so that volume and surface viewing are linked
 - ★ Click in AFNI or SUMA to change focus point, and the other program jumps to that location at the same time
 - ★ Functional overlay in AFNI can be sent to SUMA for simultaneous display
- And much more stayed tuned for the SUMA talks to come!

SUMA Teaser Movie



Color from AFNI, Images from SUMA Images captured with the 'R' recorder function, then saved as animation with Save:aGif control

Other Educational Presentations

- How to get images into AFNI or NIfTI format (program to3d)
- Detailed hands-on with using AFNI for data viewing (fun)
- Signal modeling & analysis: theory & hands-on (3dDeconvolve)
- Image registration (3dvolreg)
- Volume rendering hands-on (fun level=high)
- ROI drawing hands-on (fun level=extreme)
- Transformation to Talairach hands-on (fun level=low)
- Group analysis: theory and hands-on (3danovax)
- SUMA hands-on (fun level=pretty OK)
- Surface-based analysis
- AFNI "Jewel Box": utility programs and plugins
 - ⋆ Plus, AFNI "Jazzercise"!
- A tour of the AFNI Website

Ongoing AFNI+SUMA Projects

- Complex ANOVA models for group analyses
 - ★ Unbalanced designs, missing data, continuous covariates, multi-nested designs,
- Diffusion 'tensor' analyses
 - ★ Computation of **D** tensor, FA, tracts,
- Modeling of Gd-DTPA leakage rates into damaged tissue
- Finishing the changes to <u>3dDeconvolve</u> to simplify deconvolution analysis on paradigms with irregular timing
 - ★ And to incorporate physiological noise cancellation, and correction for EPI time series autocorrelation, and ...
- More surface-based analysis tools
- Automatic Talairach-ization + skull stripping (now working)
- Better ways to organize collections of datasets